# CENTER FOR DRUG EVALUATION AND RESEARCH

# **APPLICATION NUMBER:**

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APPROVED LABELING

# APPROV

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**AMOXIL®** 

amoxicillin

capsules, tablets, chewable tablets, and powder for oral suspension

PRESCRIBING INFORMATION 9416802

AM:L18A

#### DESCRIPTION

Amoxil formulations contain amoxicillin, a semisynthetic antibiotic, an analog of ampicillin, with a broad spectrum of bactericidal activity against many gram-positive and gram-negative microorganisms. Dhemically it is (25.5/f.6/f.)—6/f/9-1-2-amino-24-phydroxyphenyllacetamido)-3,3-dimethyl-7-oxo-4-thia-1-azabicycloj.2-0(heptane-2-carboxylic acid trihydrate. It may be represented structurally as:

The amoxicillin molecular formula is C<sub>16</sub>H<sub>19</sub>N<sub>2</sub>O<sub>4</sub>S+3H<sub>2</sub>O, and the molecular weight is 419 45

Amoxil capsules, tablets, and powder for oral suspension are intended for oral

Capsules: Each Amoxil capsule, with royal blue opaque cap and pink opaque body, contains 250 mg or 500 mg amoxicillin as the trihydrate. The cap and body of the 250-mg capsule are imprinted with the product name AMOXIL and 500 that cap and body of the 500-mg capsule are imprinted with AMOXIL and 500 inactive ingredients: D&C Red No. 28, FD&C Blue No. 1, FD&C Red No. 40, gelatin, magnesium stearate, and litanium dioxide

gearin, integression is searche, and internum double.

Tabletis: Each film-coated, capsule-shaped, pink tablet is debossed with AMOXII, centered over 500 or 875, respectively. The 875-mg tablet is scored on the reverse side linactive ingredients: colloidal solicon dioxide, crospovidone, FO&C Red No. 30 aluminum lake, hydroxypropyl methylceflulose, magnesium stearate, microcrystalline cellulose, polyethylene glycol, sodium starch glycolate, and thating dependent.

trainium dioxide. Chevrable Tablets: Each cherry-banana-peppermint-flavored tablet contains 125 mg, 200 mg, 250 mg or 400 mg amoucullin as the trihydrate. The 125-mg and 250-mg pink, void tablets are imprinted with the product name AMOXII. on one side and 125 or 250 on the other side linactive ingredients; citir, card, corn starch, FD&C Red No. 40, flavorings, glycine, mannitol, magnesium stearate, saccharin sodium, silica gel, and sucrose. Each 125-mg chewable tablet contains 0.0037 mEq (0.045 mg) of sodium. the 250-mg chewable tablet contains 0.0037 mEq (0.085 mg) of sodium.

Each 200-mg chewable tablet contains 0 0005 mEq (0 0107 mg) of sodium, the 400-mg chewable tablet contains 0.0009 mfq (0.0215 mg) of sodium. The 200 mg and 400-mg pale pink round tablets are imprinted with the product name AMOXII, and 200 or 400 along the edge of one side. Inactive ingredients aspartame 

• crospovidore NF, FD&C Red No. 40 aliminum take, flavorings, magnesium stearate and mannitol

#### • See PRECAUTIONS

Powder for Oral Suspension: Each 5 mL of reconstituted suspension contains 125 mg, 200 mg, 250 mg or 400 mg amoucillun as the trithydrate. Each 5 mL of the 125-mg reconstituted suspension contains 0.11 mEq (2.51 mg) of sodium, each 5 mL of the 250-mg reconstituted suspension contains 0.15 mEq (3.36 mg) of sodium. Each 5 mL of the 200-mg reconstituted suspension contains 0.15 mEq (3.39 mg) of sodium, each 5 mL of the 200-mg reconstituted suspension contains 0.15 mEq (4.33 mg) of sodium.

Pediatric Drops for Oral Suspension: Each ml. of reconstituted suspension contains 50 mg amoxicillin as the trithydrate and 0.03 mEg (0.69 mg) of sodium.

Amoxicillin trihydrate for oral suspension 125 mg/s ml, reconstituted) is a strawberry-flavored pink suspension, the 200 mg/s ml, 250 mg/s ml, or 50 mg/ml,) and 400 mg/s ml, are bubble-gum-flavored pink suspensions fractive ingredients: FD&C fled No. 3, flavorings, silica gel, sodium berzoate, sodium citrate, sucrose, and xanthan gum

### CLINICAL PHARMACOLOGY

CLINICAL PHARMACOLOGY
Amountain is stable in the presence of gastric acid and is rapidly absorbed after oral administration. The effect of food on the absorption of amountained inform Amount lables and Amount is suspension has been partially investigated. The 400-mg and 875-mg formulations have been studied only when administered at the start of a light meal. However, food effect studies have not been performed with the 200-mg and 500-mg formulations. Amountain diffuses readily into most body tissues and fluids, with the exception of brain and spinal fluid, except when meninges are inflamed. The half-life of amountains is 30.3 minutes. Most of the amountains excreted unchanged in the urine; its excretion can be delayed by concurrent administration of probeneous. In blood serum, amountain is as some present and in the property of the p serum, amoxicillin is approximately 20% protein-bound.

Orally administered doses of 250 mg and 500 mg amoxicillin capsules result in average peak blood levels 1 to 2 hours after administration in the range of 3.5 µg/ml, to 5.0 µg/ml, and 5.5 µg/ml, to 7.5 µg/ml, respectively.

Mean amoutcillin pharmacokinetic parameters from an open, two-part, single-dose crossover bioequivalence study in 27 adults comparing 875 mg of August (amoucillin) with 875 mg of Augustina (amoucillin) with 875 mg of Augustina (amoucillin) conditions of 35.4 ±8.1 μg/m/m, and a C<sub>max</sub> of 13.8 ±4.1 μg/m/m. Dosing was at the start of a light meal following an overnight fast.

Amoxicilin chewable tablets, 125 mg and 250 mg, produced blood levels similar to those achieved with the corresponding doses of amoxicillin oral suspensions Orally administered doses of amoxicillin suspension, 125 mg/5 ml, and 250 mg/5 ml, result in average peak blood levels 1 to 2 hours after administration in the range of 1.5 µg/ml, to 3.0 µg/ml, and 3.5 µg/ml, to 50 µg/mL, respectively

Oral administration of single doses of 400-mg *Amoxil* chewable tablets and 400-mg/5 mL suspension to 24 adult volunteers yielded comparable pharma-

Dose†	AUCo (ug.hr./ml.)	C (µg/mL) <sup>a</sup>
amoxicillin	amoxicillin (±S.D.)	amoxicallin (±S D )
400 mg (5 mL of suspension)	17.1 (3.1)	5.92 (1.62)
400 mg (one chewable tablet)	17.9 (2.4)	5.18 (1.64)

 Administered at the start of a light meal.
 Mean values of 24 normal volunteers. Peak concentrations occurred appromately 1 hour after the dose.

Detectable serum levels are observed up to 8 hours after an orally administer dose of amoxicillin. Following a 1-gram dose and utilizing a special skin w dow technique to determine levels of the artibiotic, it was noted that the peutic levels were found in the interstitial fluid. Approximately 60% of or administered dose of amoxicillin is excreted in the urine within 6 to 8 hours.

Microbology

Amoxicilin is similar to ampicillin in its bactericidal action against suscepti
organisms during the stage of active multiplication. It acts through the inittion of biosynthesis of cell wall mucopeptide. Amoxicillin has been shown
be active against most strains of the following microorganisms, both in n
and in clinical infections as described in the INDICATIONS AND USA:

## Aerobic gram-positive microorganisms: Enterococcus faecalis

Staphylococcus spp.1 (B-lactamase-negative strains only)

Streptococcus pneumoniae Streptococcus spp. (α:- and β-hemolytic strains only)

Staphylococci which are susceptible to amoxicitlin but resistant to met cillin/oxacillin should be considered as resistant to amoxicillin.

Aerobic gram-negative microorganisms: Escherichia coli (β-lactamase-negative strains only) Haemophilus influenzae (β-lactamase-negative strains only) Nesseria gonorrhoeae (β-lactamase-negative strains only) Proteus mirabilis (B-lactamase-negative strains only)

**Relicobacter** Helicobacter pylor

#### Susceptibility tests

Susceptibility tests
Dilution techniques: Quantitative methods are used to determine antimic bial minimum inhibitory concentrations (MICs). These MICs provide estimated the susceptibility of bacteria to antimicrobial compounds. The MICs should be determined using a standardized procedure. Standardized procedures a based on a dilution method 'froith or agail' or equivalent with standardized moculum concentrations and standardized concentrations of ampicillin poder. Ampicillin is sometimes used to predict susceptibility of Sireptococ. preumonae to amountlin, however, some intermediate strains have be shown to be susceptible to amountlin. Therefore, Streptococcus pneumoni-susceptibility should be tested using amountlin powder. The MIC valui should be interpreted according to the following criteria.

### For gram-positive aerobes:

Enterococcus

MIC (ug/mL) Interpretation Susceptible (S) ≥ 16 Resistant (R)

Staphylococcus

MtC (ug/mL)
≤ 0.25
≥ 0.5 Interpretation Susceptible (S) Resistant (R)

Streptococcus (except S pneur MIC (ug/mL)

Interpretation Susceptible (S)

> 8 Resistant (R) S. pneumon

(Amoxicitiin powder should be used to determine susceptibility)

MIC (ug/mL) ≤ 0.5 Interpretation Susceptible (S)

≥2 Resistant (R)

for gram-negative aerobes:

Enterobacteri MIC (ug/mL) Interpretation Susceptible (S) Intermediate (I)

≥32

MIC (ug/mL) Interpretation

Resistant (R)

 Staphylococci which are susceptible to amoxicillin but resistant to methi-cillin/oxacillin should be considered as resistant to amoxicillin. These interpretive standards are applicable only to broth microditution susceptibility tests using cation-adjusted Mueller-Hinton broth with 2.5%

c These interpretive standards are applicable only to broth microdilution test with Haemophilus influenzae using Haemophilus Test Medium (HTM).

A report of "Susceptible" indicates that the pathogen is likely to be inhibited if the antimicrobial compound in the blood reaches the concentrations usually achievable. A report of "Intermediate" indicates that the result should be

considered equivocal, and, if the microorganism is not fully susceptible to atternative, clinically feasible drugs, the test should be repeated. This category implies possible clinical applicability in body sites where the drug is physiologically concentrated or in situations where high dosage of drug can be used. This category also provides a buffer zone which prevents small uncontrolled technical factors from causing major discrepancies in interpretation. A report of "Resistant" indicates that the pathogen is not likely to be inhibited if the antimicrobial compound in the blood reaches the concentrations usually achievable; other therapy should be selected.

Standardized susceptibility test procedures require the use of laboratory control microorganisms to control the technical aspects of the laboratory procedures. Standard ampicillia powder should provide the following MIC

Microorganism	MIC (pg/mL
E. coli ATCC 25922	2 to 8
E. faecalis ATCC 29212	0.5 to 2
H. influenzae ATCC 49247°	2 to 8
S. aureus ATCC 29213	0 25 to 1

### Using amoxicillin to determine susceptibility

Microorganism S pneumoniae eumoniae ATCC 49619° MIC Range (µg/mL) 0 03 to 0 12

- This quality control range is applicable to only H influenzae ATCC 49247 tested by a broth microdilution procedure using HTM<sup>1</sup>
   This quality control range is applicable to only S preumoniae ATCC 49619 tested by the broth microdilution procedure using cation-adjusted Mueller-Hinton broth with 2-5% lysed horse blood.

Diffusion techniques: Quantitative methods that require measurement of zone diameters also provide reproducible estimates of the susceptibility of bacteria to antimicrobial compounds. One such standardized procedure-requires the use of standardized inoculum concentrations. This procedure uses paper disks impregnated with 10 µg ampiculini to test the susceptibility of microorganisms, except S pneumonae, to amoscullin Interpretation involves. the diameter obtained in the disk test with the MIC for

Reports from the laboratory providing results of the standard single disk susceptibility test with a 10-µg ampiculian disk should be interpreted accordmg to the following criteria

For gram-positive aerobes: Enterococcus Zone Diameter (mm) ≥17 Interpretation Susceptible (S)

Resistant (R) Interpretation Susceptible (S) Resistant (R) ater (mm)

<28

β-hemolytic streptococci
Zone Diameter (mm) Interpretation Susceptible (S) Intermediate (I) 19 to 25 Resistant (R)

**NOTE:** For streptococci (other than  $\beta$ -hemolytic streptococci and S pneumoniael, an ampicillin MIC should be determined.

S. pneumoniae

S. pneumoniae should be tested using a 1-µg oxacillin disk. Isolates with oxacillin zone sizes of ≥20 mm are susceptible to amoxicillin. An amoxicillin MtC should be determined on isolates of S. pneumoniae with oxacillin zone sizes of ≤19 mm.

### For gram-negative aerobes:

Interpretation Zone Diameter (mm) 14 to 16 ≤13 Resistant (R) H influ Interpretation Zone Diameter (mm) Susceptible (S) Intermediate (I) 19 to 21 Resistant (R)

- f. Staphylococci which are susceptible to amoxicillin but resistant to me cillin/oxacillin should be considered as resistant to amoxicillin
- These interpretive standards are applicable only to disk diffusion susce bility tests with H. influenzae using Haemophilus Test Medium (HTM)

nterpretation should be as stated above for results using dilution techniques As with standard dilution techniques, disk diffusion susceptibility test procedures require the use of laboratory control microorganisms. The 10-µg ampi-cillin disk should provide the following zone diameters in these laboratory test quality control strains:

Microorganism	Zone diameter (mm)
E. coli ATCC 25922	16 to 22
H. influenzae ATCC 49247h	13 to 21
S. aureus ATCC 25923	27 to 35
Using 1-µg oxacillin disk:	
Microorganism	Zone diameter (mm)
S. pneumoniae ATCC 49619	8 to 12

h. This quality control range is applicable to only H. influenzae ATCC 49247 tested by a disk diffusion procedure using HTM.?

instruction of the state of the

Susceptibility testing for Helicobacter pytori In vitro susceptibility testing methods and diagnostic products currently available for determining minimum inhibitory concentrations (MICs) and zone sizes have not been standardized, validated, or approved for testing H. pylori

Culture and susceptibility testing should be obtained in patients who fail triple therapy. If clarithromycin resistance is found, a non-clarithromycin-containing regimen should be used.

#### INDICATIONS AND USAGE

Amoul (amouscillin) is indicated in the treatment of infections due to susceptible (ONLY B-lactamase-negative) strains of the designated microorganisms in the oos listed helow

Infections of the ear, nose, and throat due to Streptococcus spp (α- and β-hemolytic strains only), Streptococcus pneumoniae, Staphylococcus spp., or H influenzae

Infections of the genitourinary tract due to E. coli, P. mirabilis, or E. faecalis Infections of the skin and skin structure due to Streptococcus spp  $\{\alpha\}$  and  $\beta$ -hemolytic strains only). Staphylococcus spp , or E coli

Infections of the lower respiratory tract due to Streptococcus spp.  $(\alpha)$  and  $\beta$ -hemolytic strains only), Streptococcus pneumoniae, Staphylococcus spp., or H influenzae

Gonorrhea, acute uncomplicated (ano-genital and urethral infections) due to

Therapy may be instituted prior to obtaining results from bacteriological and susceptibility studies to determine the causative organisms and their susceptibility to amoxicillin

Indicated surgical procedures should be performed

H. pylori eradication to reduce the risk of duodenal ulcer recurrence Triple therapy. Amoxil/clarithromycin/lansoprazole

Imple therapy Amoxii/clarithromycin plus lansoprazole as triple therapy, is Amoxii, in combination with clarithromycin plus lansoprazole as triple therapy, is AMDIAI, II COMPRISIONALLY WITH COMPRISION PLANS STANDARD PLANS STANDARD WITH THE COMPRISION PLANS STANDARD WITH THE COMPR ADMINISTRATION |

### Dual therapy: Amoxil/lans

Dual therapy: Amoni/lansoprazole Amoni/lansoprazole delayed-release capsules as dual therapy, is indicated for the treatment of patients with H. pylori infection and duodenal ulcer disease (active or one-year history of a duodenal ulcer) who are either allergic or intolerant to clarithromycin or in whom resistance to clarithromycin is known or suspected. (See the clarithromycin package insert, MICROBIOLOGY.) Eradication of H. pylori lasteen shown to reduce the risk of duodenal ulcer recurrence (See CLINICAL STUDIES and DOSAGE AND ADMINISTRATION.)

### CONTRAINDICATIONS

A history of allergic reaction to any of the penicillins is a contraindication.

WARNINGS
SERIOUS AND OCCASIONALLY FATAL HYPERSENSITIVITY (ANAPHYLACTIC)
REACTIONS HAVE BEEN REPORTED IN PATIENTS ON PENICLIIN THERAPY.
ALTHOUGH ANAPHYLAIS IS MORE REQUENT FOLLOWING PARENTERAL
THERAPY, IT HAS OCCURRED IN PATIENTS ON ORAL PENICLIINS. THESE
PEACTIONS ARE MORE LIKELY TO OCCUR IN INDIVIDUALS WITH A HISTORY
OF PENICLIIN HYPERSENSTIVITY AND/OR A HISTORY OF SENSITIVITY TO
MAITIPLE ALLERGENS. THERE HAVE BEEN REPORTS OF INDIVIDUALS WITH
A HISTORY OF PENICLIIN HYPERSENSTIVITY WHO HAVE EXPERIENCED
SEVERE REACTIONS WHEN TREATED WITH CEPHALOSPORINS. BEFORE
INITIATING THEPAPY WITH AND/OX CAREPUL INDIVIRY SHOULD BE MADE
CONCERNING PREVIOUS HYPERSENSITIVITY REACTIONS TO PENICLIINS.
CEPHALOSPORINS, OR OTHER ALLERGENS. IF AN ALLERGE PERACTION
OCCURS. AMOZE SHOULD BE DISCONTINUED AND APPROPRIATE THERAPY
INSTITUTED. SERIOUS ANAPHYLACTIC REACTIONS REQUIRE HAMEDITE EMERGENCY TREATMENT WITH EPINEPPHENE. DXYGER, INTRAVENOUS STEROIDS, AND AIRWAY MANAGEMENT, INCLUDING INTUBATION, SHOULD ALSO BE ADMINISTERED AS INDICATED. SERIOUS AND OCCASIONALLY FATAL HYPERSENSITIVITY (ANAPHYLACTIC)

Pseudomembranous colitis has been reported with nearly all antibac-terial agents, including amoxicilitia, and may range in severity from mild to life-threatening. Therefore, it is important to consider this diagnosis in patients who present with diarrhea subsequent to the administrain patients who present wit tion of antibacterial agents.

Treatment with antibacterial agents alters the normal flora of the colon and may permit overgrowth of clostridia. Studies indicate that a toxin produced by Clostridium difficile is a primary cause of "antibiotic-associated colitis."

After the diagnosis of pseudomembranous colitis has been established, appropriate therapeutic measures should be initiated. Mild cases of pseudomembranous colitis usually respond to drug discontinuation alone. In moderate to severe cases, consideration should be given to management with fluids and electrolytes, protein supplementation, and treatment with an antibacterial drug clinically effective against Clostridium difficile colitis.

### PRECAUTIONS

PRICLAUTIONS:
General: The possibility of superinfections with mycotic or bacterial pathogens should be kept in mind during therapy. If superinfections occur, amoxicillin should be discontinued and appropriate therapy instituted.

Phenyfluetonurics: Each 200 mg Amoul chewable tablet contains 1.82 mg phenyfalanine; each 400 mg chewable tablet contains 3.64 mg phenyfalanine. The Amoul suspensions do not contain phenyfalanine and can be used by phenylketonurics

Laboratory Tests: As with any potent drug, periodic assessment of renal, hepatic, and hematopoietic function should be made during prolonged therapy

All patients with gonorrhea should have a serologic test for syphilis at the time of diagnosis. Patients treated with amoxicillin should have a follow-up serologic test for syphilis after 3 months.

Oray Interactions: Probenecid decreases the renal tubular secretion of amoxicillin. Concurrent use of amoxicillin and probenecid may result in increased and prolonged blood levels of amoxicillin.

Chloramphenicol, macrolides, sulfonamides, and tetracyclines may interfere with the bactericidal effects of penicillin. This has been demonstrated in vitro, however, the clinical significance of this interaction is not well documented.

Drug/Laboratory Test Interactions: High unne concentrations of ampicillin Drug/Laboratory Test Interactions: High unne concentrations of amplicition may result in false-positive reactions when testing for the presence of glucose in urine using Clinitest®, Benedict's Solation or Fehling's Solution. Since this effect may also occur with amoxicillin, it is recommended that glucose tests based on enzymatic glucose oxidase reactions fouch as Clinistic® or Testing of the control of the

Following administration of ampiculin to pregnant women, a transient decrease in plasma concentration of total conjugated estriol, estriolglucuronide, conjugated estrone, and estradiol has been noted. This effect may also occur with amoxicilling

Carcinogenesis, Mutagenesis, Impairment of Fertility: Lovg-term stud-ies in animals have not been performed to evaluate carcinogenic potential Studies to detect mutagenic potential of amoxicillin alone have not been con-ducted, however, the following information is available from tests on a 4 1 ducted, however, the following information is available from tests on a finitude of amountallin and potassium chavilianal chaviliana chavi Augmentin was negative in the mouse micronucleus test, and in the dominant lethal assay in mice. Potassium clavulanate alone was tested in the Ames better a assay in time. Procession to continuous account our ambient assay in the mouse micronucleus test, and was negative in each of these assays. In a multi-generation reproduction study in rats, no impairment of tertility or other adverse reproductive effects were seen at doses up to 500 mg/kg (approximately 3 times the human dose in mg/m²).

Pregnancy: Teratogenic Effects. Pregnancy Category B. Reproduction studies have been performed in mice and rats at doses up to ten (10) times the human dose and have revealed no evidence of impaired fertility or harm to the numan dose and have revealed in evidence in implement returnly of failth in fetus due to amoxicillin. There are, however, no adequate and well-controlled studies in pregnant women. Because animal reproduction studies are not always predictive of human response, this drug should be used during pregnancy only if clearly needed.

Labor and Delivery: Oral ampicillin-class antibiotics are poorly absorbed during labor. Studies in guinea pigs showed that intravenous administration of ampicillin slightly decreased the uterine tone and frequency of contractions. However, it so not known whether use of amoxicillin in humans during labor or delivery has immediate or delayed adverse effects on the fetus, prolongs the duration of labor, or increases the fikelihood that forcess delivery or other obstetrical intervention or resuscitation of the newborn will be necessary.

Nursing Mothers: Penicillins have been shown to be excreted in human milk. Amoxicilin use by nursing mothers may lead to sensitization of infants. Caution should be exercised when amoxicillin is administered to a nursing

Pediatric Use: Because of incompletely developed renal function in neonates and young infants, the elimination of amountillin may be delayed. Dosing of Amouil (amountillin) should be modified in pediatric patients 12 weeks or younger (≤3 months). (See DOSAGE AND ADMINISTRATION-Neonates and infants.)

### ADVERSE REACTIONS

AUVENCE HEACTIONS

As with other penicillins, it may be expected that untoward reactions will be essentially limited to sensitivity phenomena. They are more likely to occur in individuals who have previously demonstrated hypersensitivity to penicillin and in those with a history of allergy, asthma, hay lever, or urticaria. The following adverse reactions have been reported as associated with the use of the control of the penicilling and the p nonicities

Gastrointestinal nausea, vomiting, diarrhea, and hemorrhagic/pseudomen

embranous colitis symptoms may occur during or aftantibiotic treatment. (See WARNINGS.)

Hypersensitivity Reactions: Serum sickness like reactions, erythematous mac Innanular rashes, erythema multiforme, Stevens-Johnson Syndrome, exfoltive dermatitis, toxic epidermal necrolysis, hypersensitivity vasculitis and ur caria have been reported.

NOTE: These hypersensitivity reactions may be controlled with antihistamir and, if necessary, systemic contiousteroids. Whenever such reactions ox amoxicillin should be discontinued unless, in the opinion of the physician,

condition being treated is life-threatening and amenable only to amoxicitiin

rer: A moderate rise in AST (SGOT) and/or ALT (SGPT) has been noted, but ..e significance of this finding is unknown. Hepatic dysfunction including cholestatic jaundice, hepatic cholestasis and acute cytolytic hepatitis have

Hemic and Lymphatic Systems: Anemia, including hemolytic anemia, thrombo-cytopenia, thrombocytopenic purpura, eosinophilia, leukopenia, and agranulo-cytosis have been reported during therapy with penicillins. These reactions are usually reversible on discontinuation of therapy and are believed to be hyper-sensitivity phenomena.

Central Nervous System: Reversible hyperactivity, agitation, anxiety, insomnia, confusion, convulsions, behavioral changes, and/or dizziness have been report-

Combination therapy with clarithromycin and lansourazole in clinical trials using combination therapy with amoucillin plus clarithromycin and lansoprazole, and amoucillin plus lansoprazole, no adverse reactions peculiar to these drug combinations were observed. Adverse reactions that have occurred have been limited to those that had been previously reported cicillin, clarithromycin, or lansoprazole.

<u>Irigle therapy</u>, amoxicillin/clarithromycin/lansoorazole

The most frequently reported adverse events for patients who received triple therapy were diarrhea (7%), headache (6%), and taste perversion (5%). No treatment emergent adverse events were observed at significantly higher rates with triple therapy than with any dual therapy regimen.

Dual therapy: amoxicullin/lansoprazole
The most frequently reported adverse events for patients who received amoxicillin t.i.d. plus lansoprazole t.i.d. dual therapy were diamthea (8%) and headache (7%). No treatment-emergent adverse events were observed at significantly higher rates with amoxicillin t.i.d. plus lansoprazole t.i.d. dual therapy than with lansoprazole alone.

For more information on adverse reactions with clarithromyour or lansoprazole, refer to their package inserts, ADVERSE REACTIONS

In case of overdosage, discontinue medication, treat symptomatically, and institute supportive measures as required. If the overdosage is very recent and there is no contraindication, an attempt at emesis or other means of removal. of drug from the stomach may be performed. A prospective study of 51 pediating patients at a poison-control center suggested that overdosages of less than 250 mg/kg of amoxicilin are not associated with significant clinical symptoms and do not require gastric emptying.

Interstrial neptritis resulting in oliquric renal failure has been reported in a small number of patients after overdosage with amoxicillin. Renal impairment appears to be reversible with cessation of drug administration. High blood levis may occur more readily in patients with impaired renal function because decreased renal clearance of amoxicillin. Amoxicillin may be removed from

### JOSAGE AND ADMINISTRATION

Amonii capsules, chewable tablets and oral suspensions may be given with out regard to meals. The 400-mg suspension, 400-mg chevable tablet and the 875-mg tablet have been studied only when administered at the start of a light meal. However, food effect studies have not been performed with the 200-mg

Henestes and infants aged ≤12 weeks (≤3 months)
Due to incompletely developed renal function affecting elimination of amoxicitin in this age group, the recommended upper dose of Amoxil (amoxicillin) is
30 mg/kg/day divided q12h.

### Adults and pediatric patients >3 months

Indection	Severity*	Usual Adult Dose	Usual Dose for Children >3 months <sup>6 n</sup>
Ear/nose/throat	Mild/Moderate	500 mg every 12 hours or 250 mg every 8 hours	25 mg/kg/day in divided doses every 12 hours or 20 mg/kg/day in divided doses every 8 hours
	Severe	875 mg every 12 hours or 500 mg every 8 hours	45 mg/kg/day in divided doses every 12 hours or 40 mg/kg/day in divided doses every 8 hours
Lower respiratory tract	Mild/Moderate or Severe	875 mg every 12 hours or 500 mg every 8 hours	45 mg/kg/day in divided doses every 12 hours or 40 mg/kg/day in divided doses every 8 hours

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500 mg every 25 mg/kg/day in 12 hours or divided doses Skin/skin structure Mild/Moderate 250 mg every every 12 hours 20 mg/kg/day in divided doses every 8 hours 875 mg every 12 hours or 500 mg every 45 mg/kg/day in divided doses every 12 hours 8 hours 40 mg/kg/day in divided doses every 8 hours 500 mg every 12 hours or 25 mg/kg/day in divided doses Genitourinary tract Mild/Moderate every 12 hours 250 mg every 8 hours 20 mg/kg/day in divided doses every 8 hours 875 mg every 45 mg/kg/day in Severe 12 hours or 500 mg every every 12 hours 8 hours 40 mg/kg/day in divided doses every 8 hours 3 grams as single oral dose Prepubenal children 50 mg/kg Amoxil combined with Gonombea uncomplicated ano-genital and urethral infections 25 mg/kg probenecid as a single dose. NOTE: SINCE PROBENECID IS n males and **females** CONTRAINDICATED IN CHILDREN UNDER 2 YEARS DO NOT USE THIS REGIMEN IN THESE

- Dosing for infections caused by less susceptible organisms should follow the recommendations for severe infections
- The children's dosage is intended for individuals whose weight is less than 40 kg. Children weighing 40 kg or more should be dosed according to the adult recommendations.
- Each strength of Amoxil suspension is available as a chewable tablet for use

After reconstitution, the required amount of suspension should be placed After reconstitution, the required amount of suspension should be packed directly on the child's tongue for swallowing. Alternate means of administra-tion are to add the required amount of suspension to formula, milk, fruit juice, water, ginger ale, or cold drinks. These preparations should then be taken immediately To be certain the child is receiving full dosage, such preparations

All patients with gonorrhea should be evaluated for syphilis. (See PRECAU-TIONS - Laboratory Tests.)

Larger doses may be required for stubborn or severe infections

General: It should be recognized that in the treatment of chronic urinary tract infections, frequent bacteriological and clinical appraisals are necessary. Smaller doses than those recommended above should not be used. Even higher doses may be needed at times. In stubborn infections, therapy may be er ooses may be needed at times. In stubbom infections, therapy may be required for several weeks. It may be necessary to continue clinical and/or bacteriological follow-up for several months after cessation of therapy. Except for gonorrhea, treatment should be continued for a minimum of 48 to 72 hours beyond the time that the patient becomes asymptomatic or evidence of bactrial eradication has been obtained. It is recommended that there be at least 10 days' treatment for any infection caused by Streptococcus pyogenes to prevent the occurrence of acute rheumatic fever.

H. pylori eradication to reduce the risk of duodenal ulcer recurrence liriple therapy: Amosil/clarithromycir/lansoprazole
The recommended adult oral dose is 1 gram Amosil, 500 mg clarithromycin, and 30 mg lansoprazole, all given twice daily (q12h) for 14 days. (See INDICATIONS AND USAGE.)

Dual therapy: Amoxil/lansoprazole
The recommended adult oral dose is 1 gram Amoxil (amoxicifin) and 30 mg lansoprazole, each given three times daily (q8h) for 14 days. [See INDICATIONS AND USAGE.]

Please refer to clarithromycin and lansoprazole full prescribing information for CONTRAINDICATIONS and WARNINGS, and for information regarding dosing in elderly and renally impaired patients.

Dosing recommendations for adults with impaired renal function: Patients with impaired renal function do not generally require a reduction in dose unless the impairment is severe Severely impaired patients with a glome

should receive 500 mg or 250 mg every 24 hours, depending on severity of the

Hemodialysis patients should receive 500 mg or 250 mg every 24 hours, depending on severity of the infection. They should receive an additional dose both during and at the end of dialysis.

There are currently no dosing recomwith impaired renal function. endations for pediatric patients

Directions For Mixing Oral Suspension
Prepare suspension at time of dispensing as follows: Tap bottle until all powder
flows freely. Add approximately 1/3, of the total amount of water for reconstitution (see table below) and shake vigorously to wet powder. Add remainder of
the water and again shake vigorously.

	125 mg/5 mL	Amount of Water
Bottle Size	•	Required for Reconstitution
80 mL		62 mL
100 mL		78 mL
150 mL		116 mL
Each teaspoonful (5 mL) v	will contain 125 mg a	moxicillin.

	200 mg/5 mL	Amount of Water
Bottle Size	•	Required for Reconstitution
5 mt		5 mL
50 mL		39 mL
75 mL		57 mL
100 mL		76 mL
Each teaspoonful (5 m	L) will contain 200 mg a:	moxicillin

Amount of Water 250 ma/5 ml. **Bottle Size** Required for Reconstitution

8U IIIL		33 110
100 mL		74 mL
150 mL		111 mL
Each teaspoonful (5 mL) v	vill contain 250 mg ar	noxicillin.
	400 mg/5 mL	Amount of Water
Bottle Size	-	Required for Reconstitution
5 mL		5 mL
SO ml		36 ml

75 ml 54 mt 100 mL Each teaspoonful (5 mt.) will contain 400 mg amoxicillin

**Directions For Mixing Pediatric Drops** 

Prepare pediatric drops at time of dispensing as follows. Add the required amount of water (see table below) to the bottle and shake vigorously. Each ml. of suspension will then contain amoxicillin trihydrate equivalent to 50 mg.

	Amount of Water
Bottle Size	Required for Reconstitution
15 mL	12 mL
30 mL	23 mL

NOTE: SHAKE BOTH ORAL SUSPENSION AND PEDIATRIC DROPS WELL BEFORE USING. Keep bottle tightly closed. Any unused portion of the reconstituted suspension must be discarded after 14 days. Refrigeration preferable, but not required.

HOW SUPPLIED Amoxii (amoxicillin) Capsules Each capsule contains 250 mg or 500 mg amoxicillin as the trihydrate.

	The last cobserve	
NDC 0029-6006-30		bottles of 100
NDC 0029-6006-32		bottles of 500
	500-mg Capsule	
NDC 0029-6007-30	• •	bottles of 100
NDC 0029-6007-32		bottles of 500

oxicillia) Tablets. Each tablet contains 500 mg or 875 mg amoxicillin as the trihydrate.

300-si	al refiles
NDC 0029-6046-12	bottles of 20
NDC 0029-6046-20	bottles of 100
NDC 0029-6046-25	bottles of 500
875-a	ng Tablet
NDC 0029-6047-12	bottles of 20
NDC 0029-6047-20	bottles of 100
NDC 0029-6047-25	bottles of 500

Amoxici (amoxicittin) Chewable Tablets. Each cherry-banana-peppermint-flavored tablet contains 125 mg, 200 mg, 250 mg or 400 mg amoxicillin as the

	125-mg Tablet	
NDC 0029-6004-39		bottles of 60
	200-mg Tablet	
NDC 0029-6044-12	-	bottles of 20
NDC 0029-6044-20		bottles of 100
	250-mg Tablet	
NDC 0029-6005-13	•	bottles of 30
NDC 0029-6005-30		bottles of 100

rerity of the 24 hours.

c pat

NDC 0029-8045-12 NDC 0029-6045-20

bottles of 20 hottles of 100

80-mL bottle 100-mL bottle 150-mL bottle

50-mi. bottle 75-mi. bottle 100-mi. bottle

80-ml hottle

50-mL bottle 75-mL bottle 100-mL bottle

30-ml. bottle

290-mg unit dose bottle 400-mg unit dose bottle

Amoriil (emexicitie) for Oral Suspension. Each 5 mt. of reconstituted strawberny-flavored suspension contains 125 mg amoricitiin as the trihydrate. Each 5 mt. of reconstituted bubble-grun-flavored suspension contains 200, 250, or 400 mg amoricifiin as the trihydrate.

125 mg/5 mL

200 mg/5 mL

250 mg/5 mL

400 mg/5 mL

400-mg Tablet

all powder reconstitu- mainder of
dae

NOC 0025-0000-22	
NDC 0029-6048-54 NDC 0029-6048-55 NDC 0029-6048-58	

NDC 0029-6008-21

NOC 0029-6008-23

NDC 0029-6009-21 NDC 0029-6009-23

NDC 0029-6049-54 NDC 0029-6049-55 NDC 0029-6049-59 NDC 0029-6048-18 NDC 0029-6049-18

Amoxid (amoxicitlin) Pediatric Drops for Oral Suspension. Each ml. of bubble-gum-flavored reconstituted suspension contains 50 mg amoxicillin as the tribydrate

NEDC 0029-6035-20 NDC 0029-6038-39

Store capsules, unreconstituted powder, and 125-mg and 250-mg chewable tablets at or below 20°C (66°T). The 200-mg and 400-mg chewable tablets may be stored at or below 25°C (77°F) Store 500-mg and 875-mg tablets at or below 25°C (77°F) Dispense in a right container

CLUNICAL STUDIES

Randomized, double-blind clinical studies performed in the U.S. in patients with H pydori and doudenal ulcer is studies performed in the U.S. in patients with H pydori and doudenal ulcer disease (defined as an active ulcer or history of an ulcer within one year) evaluated the efficacy of lansoprizable in combination with amouncillin capsules and clarithromycin tablets as triple 14-day therapy, or in combination with amouncillin capsules as dual 14-day therapy, or in combination with amouncillin capsules as dual 14-day therapy. for the eradication of *H. pylon*. Based on the results of these studies, the safety and efficacy of two different eradication regimens were established

Triple therapy: amoxicillin 1 gram b i d /clarithromycin 500 mg b i d /lansoprazole 30 mg b i d

Dual therapy: amoxicillin 1 gram t i d./lansoprazole 30 mg t.i d

All treatments were for 14 days *H pylori* eradication was defined as two agative tests (culture and histology) at 4 to 6 weeks following the end of atment.

.riple therapy was shown to be more effective than all possible dual therapy combinations. Dual therapy was shown to be more effective than both monotherapies. Eradication of *H. pylon* has been shown to reduce the risk of

H. pylori Eradication Rates – Triple Therapy (amoxicillin/clarithromycin/lansoprazole) Percent of Patients Cured [95% Confidence Interval] (Number of Patients)

<b>(</b>			
Study	Triple Therapy  Evaluable Analysis <sup>†</sup>	Triple Therapy	
		Intent-to-Treat Analysis*	
Study 1	92° (80.0–97.7] (n=48)	86° [73.3–93.5] (n=55)	
Study 2	86° [75.7–93.6] (n=66)	83" [72.0–90.8] (n=70)	

rtution

'S WEII

500 mg

- This analysis was based on evaluable patients with confirmed duodenal ulcer (active or within one year) and *H. pylori* infection at baseline defined as at least two of three positive endoscopic tests from Q.Otest\*. (Detta West Ltd., Bentley, Australia), histology and/or cutture. Patients were included in the analysis if they completed the study. Additionally, if patients dropped out of the study due to an adverse event related to the study drug, they were included in the analysis as failures of therapy. Patients were included in the analysis if they had documented *H. pylori* infection at baseline as defined above and had a confirmed duodenat ulcer (active or within one year). All dropouts were included as failures of therapy.
- apy.
  (p<0.05) versus lansoprazole/amoxicillin and lansoprazole/clarithromycin
- dual therapy (p<0.05) versus clarithromycin/amoxicillin dual therapy

Study	Dual Therapy Evaluable Analysis <sup>1</sup>	Dual Therapy
		Intent-to-Treat Analysis**
Study 1	77" [62.5-87.2] [n=51]	70** [56.8–81.2] (n=60)
Study 2	66 <sup>55</sup> [51.9-77.5] (n=58)	61 <sup>54</sup> [48.5–72.9] (n=67)

- This analysis was based on evaluable patients with confirmed duodenal ulcer (active or within one year) and *H. pylori* infection at baseline defined as at least two of three positive endoscopic tests from CLOtest\*\*, histology and/or culture. Patients were included in the analysis of they completed the study. Additionally, if patients dropped out of the study due to an adverse event related to the study drug, they were included in the analysis as failures of therapy.

  11 Patients were included in the analysis if they had documented *H. pylori* infection at baseline as defined above and had a confirmed duodenal ulcer (active or writhin one year). All dropouts were included as failures of therapy.

- apy.

  ‡‡ (p<0.05) versus lansoprazole alone.

  §§ (p<0.05) versus lansoprazole alone or amoxicillin alone

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